(PCT Article 36 and Rule 70)

27 APR 2005

			N- (II- reference			was a second second		
Applicant's or agent's file reference PRD2009-PCTf				FOR FURTHER ACTION  See Notification of Transmittal of International Preliminary Examination Report (Form PCT/PEA/416)				
International application No.				International filing date (d	ay/month/year)	Priority date (day/month/year) 31.10.2002		
PCT/EP 03/11792				23.10.2003		31.10.2002		
International Patent Classification (IPC) or both national classification and IPC								
CIZU	C12Q1/68							
Applicant LANGUETICA NIV et al.								
JANSSEN PHARMACEUTICA N.V. et al.								
This international preliminary examination report has been prepared by this International Preliminary Examining     Authority and is transmitted to the applicant according to Article 36.								
2. T	his F	REPO	ORT consists of a total of	of 5 sheets, including th	s cover shee	ı.		
	☐ This report is also accompanied by ANNEXES, i.e. sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).							
 	These	ann	exes consist of a total of	of sheets.				
<u> </u>	•							
_				tata a ka ka a fallassina ika				
3.	This r	epor	t contains indications re	elating to the following ite	ems:	•		
1			Basis of the opinion					
1			Priority	and the second decision		ve etan and industrial applicability		
1					veity, inventi	ve step and industrial applicability		
1		⊠ ⊠	Lack of unity of invent		h regard to n	ovelty, inventive step or industrial applicability;		
. '	V		citations and explanat	ions supporting such sta	tement			
,	۷I		Certain documents cit	ed				
1	VII   Certain defects in the in				nternational application			
'	VIII		Certain observations	on the international appl	cation			
Date of submission of the demand Date of completion of this report								
Date of submission of the demand					Date of comp	·		
25.03.2004					10.01.200	5		
Name and mailing address of the International preliminary examining authority:				nal	Authorized O	fficer		
Premu		Eu	ropean Patent Office 80298 Munich		Grankant			
	<u>0)</u> ))	Tel	. +49 89 2399 - 0 Tx: 5236	656 epmu d	Grosskopf			
		Fa	x: +49 89 2399 - 4465		Telephone N	D. +49 89 2399-8714		

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<ol> <li>Basis of the rep</li> </ol>	oort	i
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1. With regard to the **elements** of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)):

	Des	cription, Pages					
	1-53	3	as originally filed				
	Seq	uence listings part o	f the description, Pages				
	54-6	64	as originally filed				
	Clai	ms, Numbers					
	1-31	l	as originally filed				
	Dra	wings, Sheets					
	1/13	-13/13	as originally filed				
2.	With lang	With regard to the <b>language</b> , all the elements marked above were available or furnished to this Authority in the anguage in which the international application was filed, unless otherwise indicated under this item.					
	These elements were available or furnished to this Authority in the following language: , which is:						
		the language of a tran	nslation furnished for the purposes of the international search (under Rule 23.1(b)).				
		the language of public	cation of the international application (under Rule 48.3(b)).				
		the language of a trar Rule 55.2 and/or 55.3	nslation furnished for the purposes of international preliminary examination (under ).				
3.	With inte	n regard to any <b>nucleo</b> rnational preliminary e	otide and/or amino acid sequence disclosed in the international application, the xamination was carried out on the basis of the sequence listing:				
	×	contained in the inter	national application in written form.				
	$\boxtimes$	filed together with the	international application in computer readable form.				
		furnished subsequent	tly to this Authority in written form.				
	☐ furnished subsequently to this Authority in computer readable form.						
		The statement that the in the international ap	e subsequently furnished written sequence listing does not go beyond the disclosure oplication as filed has been furnished.				
		The statement that the listing has been furnish	e information recorded in computer readable form is identical to the written sequence shed.				
4.	The	amendments have re	sulted in the cancellation of:				
		the description,	pages:				
		the claims,	Nos.:				
		the drawings,	sheets:				

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5.  This report has been established as if (some of) the amendments had not been made been considered to go beyond the disclosure as filed (Rule 70.2(c)).								
		(Any replacement sheet conta report.)	ining s	euch amendn	nents must be referred to under item 1 and annexed to this			
6.	Add	Additional observations, if necessary:						
IV	. Lac	k of unity of invention						
1.	In re	esponse to the invitation to rest	al fees, the applicant has:					
		restricted the claims.						
☐ paid additional fees.								
		paid additional fees under prof	fees under protest.					
		☐ neither restricted nor paid additional fees.						
2.		This Authority found that the requirement of unity of invention is not complied with and chose, according to Rule 68.1, not to invite the applicant to restrict or pay additional fees.						
3.	This	This Authority considers that the requirement of unity of invention in accordance with Rules 13.1, 13.2 and 13.3 is						
□ complied with.								
☐ not complied with for the following reasons:								
	see	see separate sheet						
4.		Consequently, the following parts of the international application were the subject of international preliminary examination in establishing this report:						
☐ all parts.								
	☒	the parts relating to claims No	claims Nos. 1-5,26-31 .					
V.		easoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; itations and explanations supporting such statement						
1.	Stat	ement						
	Nov	elty (N)	Yes: No:	Claims Claims	1-5 26-31			
	Inve	entive step (IS)	Yes: No:	Claims Claims	1-5			
	Indu	strial applicability (IA)	Yes: No:	Claims Claims	1-5, 26-31			

2. Citations and explanations

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see separate sheet



# **EXAMINATION REPORT - SEPARATE SHEET**

#### Ad item IV and V:

This Authority is in complete agreement with the opinion of the Search Authority as far as unity is concerned for the reasons outlined in the search report.

Since no additional search fees have been aid this opinion will be restricted to Claims 1 to 5 and 26 to 31 as far as SEQ ID NO: 1 is concerned.

The nucleotide sequence according to SEQ ID NO: 1 is a known sequence (see D1; DATABASE EMBL [Online] 16 July 1999 (1999-07-16), XP002249222 retrieved from EBI Database accession no. Al842377). As a consequence, the products according to claims 26 to 31 lack novelty and/or are devoid of any inventive merit.

The expression of the known sequence has been found to be responsive to the corticotrophin releasing hormone (CRH) in the CNS. Such sequences are well known in the art (see the other documents cited in the search report).

Thus, the identification of (further) genes which are responsive to CRH and their obvious applications, respectively merely the measurement of their level of transcription (see Claims 1 to 5) must be considered as being devoid of any inventive merit.

An inventive activity could at best be acknowledged for the use of said known sequence e.g. in diagnosis or therapy. In this respect, however, the application fails to provide any evidence or examples for any of the claimed sequence, let alone for SEQ ID NO: 1. For the assessment of the present claims 1 to 5 on the question whether they are industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.